RESEARCH COMMUNICATION

Homologous recombination as a mechanism to recognize repetitive DNA sequences in an RNAi pathway

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Quelling is an RNAi-related phenomenon that post-transcriptionally silences repetitive DNA and transposons in Neurospora. We previously identified a type of DNA damage-induced small RNA called qiRNA that originates from ribosomal DNA. To understand how small RNAs are generated from repetitive DNA, we carried out a genetic screen to identify genes required for qiRNA biogenesis. Factors directly involved in homologous recombination (HR) and chromatin remodeling factors required for HR are essential for qiRNA production. HR is also required for quelling, and quelling is also the result of DNA damage, indicating that quelling and qiRNA production share a common mechanism. Together, our results suggest that DNA damage-triggered HR-based recombination allows the RNAi pathway to recognize repetitive DNA to produce small RNA.

Supplemental material is available for this article.

Received November 2, 2012; revised version accepted December 21, 2012.

RNAi is a mechanism of gene silencing conserved from fungi to mammals [Catalanotto et al. 2006; Buchler and Moazed 2007; Ghildiyal and Zamore 2009]. Most of the RNAi-related pathways are initiated by production of dsRNA, which is cleaved by Dicer to generate siRNAs. The siRNAs are loaded into a complex containing an Argonaute (Ago) family protein to mediate gene silencing of homologous RNAs. RNAi is an ancient genome defense mechanism that silences transposons and viral infection [Sijen and Plasterk 2003; Siomi et al. 2008; Wang et al. 2010; Chang et al. 2012]. Consistent with this role of RNAi, a significant portion of eukaryotic small RNAs, including siRNA and the piwi-interacting small RNAs (piRNAs), are produced from repetitive DNA loci in fungi, plants, and animals and target active transposon sequences or their relics [Siomi et al. 2008; Ghildiyal and Zamore 2009]. How small RNAs are specifically produced from repetitive DNA loci is not clear.

Quelling in the filamentous fungus Neurospora crassa was one of first known RNAi-related phenomenon [Romano and Macino 1992; Catalanotto et al. 2006; Chang et al. 2012]. Quelling is triggered by multiple copies of transgenes and produces transgene-specific siRNA [Catalanotto et al. 2006]. In the quelling pathway, QDE-1 (Quelling-Deficient-1) first acts as a DNA-dependent RNA polymerase [DdRP] to produce aberrant RNA [aRNA] from the repetitive transgene loci and then uses its RNA-dependent RNA polymerase [RdRP] activity to convert aRNA into dsRNA [Cogoni and Macino 1999a; Lee et al. 2010]. This process requires the RecQ DNA helicase QDE-3 and the ssDNA-binding complex replication protein A [RPA] [Cogoni and Macino 1999b; Nolan et al. 2008; Lee et al. 2010]. dsRNA is then processed by Dicer proteins to produce siRNA, which is then loaded onto the Argonaute protein QDE-2 to mediate post-transcriptional gene silencing [Maiti et al. 2007]. In most Neurospora strains, the ribosomal DNA locus has the only highly repetitive DNA sequences. The potent silencing effect of quelling on repetitive transgenes suggests that it is an anti-transposon response. Consistently, the quelling pathway suppresses the replication of a functional transposon [Nolan et al. 2005].

We previously discovered that DNA damage induces the expression of QDE-2 and a class of small RNAs named qiRNAs for their association with QDE-2 [Cecere and Cogoni 2009; Lee et al. 2009, 2010]. qiRNAs originate from the rDNA locus, which contains ~200 copies of rDNA repeats, and their production depends on QDE-1, QDE-3, Dicers, and RPA. qiRNA levels are very low under normal growth conditions. DNA damage-induced small RNAs were recently also discovered in Arabidopsis, flies, and mammals [Francia et al. 2012, Michalik et al. 2012, Wei et al. 2012], suggesting that DNA damage is a common trigger for small RNA production in eukaryotes.

How does the quelling pathway specifically target repetitive DNA sequences? What is the mechanistic link between quelling and the DNA damage-induced qiRNA production? Here we show that the homologous recombination (HR) process is essential for qiRNA production and quelling. We further show that the qiRNA and the quelling-induced siRNA share the same biogenesis mechanism. Together, our results suggest that Neurospora uses HR to distinguish repetitive DNA loci from the rest of the genome.

Results and Discussion

A genetic screen identified genes required for HR to be essential for qiRNA production

Treatment of Neurospora with DNA damage agents such as histidine, hydroxyurea (HU), or ethyl methanesulfonate (EMS) induces the production of qiRNA and induction of QDE-2 protein expression [Lee et al. 2009]. In addition, the DNA damage-induced QDE-2 expression is abolished in mutants defective for qiRNA production and quelling [Lee et al. 2009, 2010]. To identify additional components of qiRNA and quelling pathways, we reasoned that mutants deficient in DNA damage-induced QDE-2 expression would also be defective in qiRNA and...
and aRNA production in a series of *Neurospora* DNA repair and checkpoint pathway mutants [Fig. 1 D,E; data not shown]. These mutants included rad18 (post-replication repair), rad5 (post-replication repair), ku70/mus-51 and ku80/mus-52 [nonhomologous end-joining], atr/mus-9 and chk2 [checkpoint], purk [single-strand break and base excision repair], and msh3 [mismatch repair]. In contrast to the HR mutants, the HU-induced qiRNA and aRNA could still be produced in these mutants. Moreover, the qde-1 and qde-3 mRNA levels in these mutants and in the HR mutants were comparable with those in the wild-type strain, indicating that the phenotype observed in the HR mutants is not due to low QDE-1 or QDE-3 levels [Supplemental Fig. S3]. These results indicate that HR, but not other DNA repair and checkpoint pathways, is required for qiRNA synthesis in a step upstream of aRNA production.

**Chromatin remodeling factors are required for qiRNA production due to their roles in HR**

In our screen, several genes encoding for ATP-dependent chromatin remodeling enzymes were also found to be required for the histidine-induced QDE-2 expression [Fig. 2A; data not shown]. These genes encode the *Neurospora* homologs of *swr1, isw1*, and *chd1* [NCU09993, NCU03875, and NCU03060, respectively]. As shown in Figure 2B, both qiRNA and aRNA production induced by HU were completely or mostly abolished in the *swr1, isw1*, and *chd1* mutants, indicating the important roles of these proteins in qiRNA biogenesis. These results further suggest that qiRNA and aRNA production is regulated at the chromatin level.

These chromatin remodeling enzymes use ATP to alter histone–DNA contacts, causing changes in the status of chromatin by moving and restructuring nucleosomes [Clapier and Cairns 2009]. Chromatin remodeling factors INO80 and SWR1 have been previously reported to be associated with DNA repair processes, and INO80 is involved in the HR process [Vignali et al. 2000; van Attikum et al. 2007]. ISW1 is involved in replication initiation and promotes replication fork progression [Vincent et al. 2008]. Because HR is required for qiRNA biogenesis, we hypothesized that these enzymes are also involved in HR. Thus, we examined the recombination rates at the *methyltryptophan resistance* (*mtr*) locus in different *Neurospora* strains by transforming cells with a construct containing the bialaphos resistance gene (*bar*) that can disrupt the *mtr* gene by HR [Ishibashi et al. 2006]. The targeting of the *bar* gene into the *mtr* gene results in transformants that are resistant to both bialaphos and the amino acid analog *p*-fluorophenylalanine [FPA]. The wild-type strain has a recombination rate of ~20% [Fig. 2C], which is a typical HR rate in *Neurospora*. As expected, the HR rates of the *ku80* strain and the *rad51* strain are nearly 100% and 0%, respectively. Consistent with a role of SRS2 in suppressing HR, the srs2 mutant exhibited an ~50% HR rate. On the other hand, the HR rates were very low or completely abolished in the *swr1, isw1*, and *chd1* mutants. These results suggest that these chromatin remodeling enzymes are required for qiRNA production due to their nonredundant roles in HR and that the chromatin remodeling factors act collaboratively to regulate chromatin status during the qiRNA production process.
The role of HR in small RNA biogenesis

The requirements for HR and the RecQ DNA helicase QDE-3 in qRNA production raised the possibility that recombination of rDNA repeats results in production of an “aberrant” DNA structure that acts as a trigger for qRNA production. To test this hypothesis, we subjected unsynchronized Neurospora cultures to HU treatment, extracted genomic DNA, and performed two-dimensional (2D) electrophoresis assays. In the absence of HU, little or no replication intermediates of rDNA could be observed [Fig. 4]. The HU treatment resulted in the accumulation of rDNA-specific replication intermediates in both the wild-type and srs2 strains, which are represented as an “arc” in the 2D electrophoresis assay (Brewer and Fangman 1987). In addition, a low level of recombination intermediates [Fig. 4, indicated by an open arrow], which are typically represented as a spike above the replication intermediates

Figure 2. ATP-dependent chromatin remodelers are involved in the qRNA pathway. (A) QDE-2 Western blot analysis of the indicated strains. Cultures were grown in histidine for 2 d. (B) Northern blot analysis showing qRNA and aRNA production in different strains after 2 d of treatment. (C) HR assays showing the HR rates of the indicated strains. n = 3, error bar indicates SD.

qRNA production requires DNA replication

HR is the predominant DNA double-strand break (DSB) repair pathway during S and G2 phases, when replicated sister chromatin provides a template for HR-dependent repair (Sancar et al. 2004). The accumulation of recombination intermediates correlates with the accumulation of replication intermediates, indicating that HR activity occurs most frequently during DNA synthesis [Zou and Rothstein 1997]. To determine whether qRNA production requires DNA replication, we treated the wild-type Neurospora with different concentrations of HU. At low concentrations of HU (0.5 and 1 mg/mL), qRNA and aRNA production were induced, whereas at 8 mg/mL HU, a concentration at which DNA replication is completely blocked [Srivastava et al. 1988], qRNA and aRNA production were abolished [Fig. 3A,B]. Similarly, treatment of wild-type cells with EMS also led to the induction of qRNA at low concentrations but a blockade of qRNA synthesis at a high concentration [Fig. 3A]. A similar HU-dose-dependent response was also observed in an atm mutant [Fig. 3B]. These results suggest that qRNA biogenesis requires DNA replication even though qRNA production is induced by modest DNA damage that can result in partial replication inhibition.

To further confirm our conclusion, we created Neurospora strains in which one of the essential components for DNA replication, proliferating cell nuclear antigen (PCNA), can be silenced by quinic acid (QA)-inducible pcna-specific dsRNA (Cheng et al. 2005). PCNA is a processivity factor that forms a complex with DNA polymerase and acts as a clamp that tethers DNA polymerase to the DNA template [Kelman 1997]. Figure 3C shows the race tube results that compared the growth phenotypes of the wild-type strain and two independent pcna knockdown strains (dpcna). In the presence of QA, the cell growth of the dpcna strains was dramatically inhibited, indicating that DNA replication was inhibited by the silencing of pcna [Fig. 3C]. In the presence of QA, the HU-induced qRNA production was completely abolished in the dpcna strains [Fig. 3D]. To examine whether the inhibition of DNA replication leads to a nonspecific loss of all small RNA, we examined the levels of an siRNA at different concentrations of HU and found that albino-1 (al-1) siRNA (produced from a double-stranded al-1 hairpin RNA) was expressed at similar levels at all concentrations of HU tested [Fig. 3E]. This result suggests that HU treatment does not affect the stability of small RNA. Together, these results suggest that qRNA biogenesis requires DNA replication and occurs during the S phase of the cell cycle.

rDNA-specific recombination intermediates accumulate upon DNA damage

The role of HR in small RNA biogenesis

Figure 3. DNA replication is required for qRNA biogenesis. (A) Northern blot analysis showing the levels of qRNA in the wild-type strain grown in the indicated concentrations of HU and EMS. (B) Northern blot analysis showing aRNA production in different strains. (C) Race tube analysis showing the growth of the indicated strains in race tubes. dpcna #1 and #2 strains are two independent dsRNA knockdown strains in which pcna dsRNA expression can be induced by QA (1 x 10⁻⁷ M). The black lines indicate cell growth fronts that were marked every 24 h. (D) Northern blot analysis showing elimination of qRNA production in the dpcna strains. Culture medium contained 10⁻⁷ M QA. (E) Northern blot analysis showing the levels of al-1 siRNA in the dsl-1 strain that expresses al-1-specific dsRNA.
Quelling requires HR and is also induced by DNA damage

Quelling-triggered gene silencing and siRNA production occur under normal growth conditions and do not require DNA damage agent treatment. However, the similarity between quelling and qiRNA biogenesis suggest that these two phenomena may be mechanistically the same. To test this hypothesis, we performed quelling assays by transforming *Neurospora* with an *al-1* transgene. As shown in Figure 5A, 29% of the wild-type transformants exhibited quelling, as indicated by the change of conidia color from orange to yellow or white. In contrast, very low quelling efficiency was observed in the *rad51*, *rad54* strains, indicating that, like QDE-3, the HR components are required for quelling.

Because repetitive DNA is known to be a major cause of genome instability due to recombination (Bzymek and Lovett 2001; Vater et al. 2011), we hypothesized that quelling is also caused by DNA damage due to repetitive transgenes. Thus, we examined whether the quelling-triggered siRNA, like qiRNA production, is triggered by DNA damage and requires DNA replication. We reasonned that a fully quelled strain might already be subject to high levels of DNA replication stress at the quelled locus and that further DNA damage treatment would not further promote the production of siRNA. Therefore, a partially *al-1*-quelled wild-type transformant (yellow conidia) was isolated and treated with HU at different concentrations. As shown in Figure 5B, the level of *al-1* siRNA was induced to a level that was similar to that of a fully quelled strain at low concentrations of HU, but the production of siRNA was completely abolished at 4 mg/mL HU. As expected, 0.5 mg/mL HU treatment of the partially quelled strain resulted in a decrease of *al-1* mRNA to a level that was comparable with that of the fully quelled strain (Supplemental Fig. S4). These results suggest that the quelling-triggered siRNA production is also the result of DNA damage and requires DNA replication.

To determine whether the HR pathway is directly involved in quelling, we expressed c-Myc-tagged RAD51 in an *al-1*-quelled strain and performed a chromatin immunoprecipitation (ChIP) assay using a monoclonal c-Myc antibody. As shown in Figure 5C, a significant enrichment of Myc-RAD51 was observed at the *al-1* transgene locus. Taken together, these results indicate that quelling and qiRNA biogenesis share the same pathway and that HR is also an essential step in quelling.

Lack of qiRNA production under normal growth conditions is due to mechanisms that prevent rDNA recombination

Why is qiRNA only induced after DNA damage treatment, whereas the quelling-triggered siRNA is produced under normal growth conditions? The eukaryotic rDNA region consists of several hundred copies of tandem rDNA repeats that each contains a potential origin of DNA replication. Because of its highly repetitive nature, hyperrecombination can occur at a rDNA locus, resulting in genome instability and chromosomal rearrangements. Previous studies have shown that several mechanisms, including rDNA transcriptional silencing and regulation of rDNA replication, prevent rDNA hyperrecombination and maintain rDNA copy numbers (Calzada et al. 2005; Huang et al. 2006). During rDNA replication, the intergenic replication fork barrier sites in rDNA repeats can
stall replication forks unidirectionally so that replication of the rDNA repeats occurs in the same direction as rDNA transcription. The stalled replication forks are protected and maintained by a fork protection complex, which prevents the collapse of the replication fork and is important for maintaining genome stability at the rDNA region. In yeast, the components of this complex have been shown to be important for the maintenance and progression of the replication fork at the rDNA replication barrier sites (Krings and Bastia 2004; Mohanty et al. 2006).

We hypothesized that mechanisms that suppress rDNA recombination during replication inhibit qiRNA production under normal growth conditions. The Neurospora swi3 (NCU01858), mrc1 (NCU04321), and mcl1 (NCU08484) genes encode the homologs of the fission yeast Swi3p, Mrclp, and Mcllp, respectively, which are part of the fork protection complex. To test our hypothesis, we examined the production of qiRNA and aRNA in the swi3, mrc1, and mcl1 mutants. As shown in Figure 5D, the levels of both qiRNA and aRNA were high even in the absence of DNA damage agent treatment in these mutant strains. In the fission yeast, Swi3 is important for the replication fork arrest in the rDNA region, and deletion of swi3 resembles the HU-induced replication stress at the rDNA region (Krings and Bastia 2004). Quelling assay results showed that these replication fork protection mutants have normal quelling efficiency (Supplemental Table S1), suggesting that these factors do not have a major impact on repetitive transgenes. These results suggest that the difference between qiRNA production and quelling is that the rDNA locus is normally actively protected from HR, but the repetitive transgene loci are not, thus, quelling occurs under normal growth conditions.

In this study, we identified HR as an essential process for the DNA damage-induced qiRNA production and quelling. Because qiRNA and quelling are both produced from repetitive DNA loci and the normally protected rDNA locus is the only highly repetitive DNA locus in the Neurospora genome, our results suggest that HR is a mechanism that can distinguish repetitive foreign DNA from the rest of the genome. Consistent with this conclusion, the quelling pathway has been shown to suppress transposon proliferation in Neurospora [Nolan et al. 2005].

Our study provides important insights into the mechanism of small RNA production from repetitive DNA loci. Even though qiRNA and the quelling-induced siRNA may appear to be triggered by different cues, we showed that the upstream mechanism for their production is the same and that both are results of DNA damage. First, the biogenesis of both types of small RNA require the same set of components, including QDE-1, QDE-3, Dicers, RPA, and the HR components. Second, like qiRNA, the transgene-induced siRNA can also be induced by DNA damage and requires DNA replication. Repetitive DNA is known to be a major source of genome instability in different organisms due to HR [Bezemek and Lovett 2001; Vader et al. 2011]. Therefore, even under normal growth conditions, repetitive transgenes lead to DNA replication stress or DSBs, resulting in the production of transgene-specific siRNA. Third, in mutants that are deficient in maintaining fork stability and progression in the rDNA locus, qiRNA levels are high without DNA damage treatment (Fig. 5D). This result suggests that the rDNA locus is normally protected to maintain its stability and suppress DNA recombination.

aRNA is the precursor of dsRNA. The production of aRNA and dsRNA is catalyzed by the DdRP/RdRP QDE-1 [Lee et al. 2010]. How does QDE-1 recognize the quelled locus and rDNA? How does DNA damage trigger the production of qiRNA and siRNA? Our genetic screening results indicate that only the HR process, but not other DNA repair or checkpoint pathways, is required for qiRNA production. After DSBs, the Rad51-coated ssDNA, with the help of Rad54, invades the sister chromatin to form recombination intermediates. Even though how repetitive DNA is recognized by the quelling machinery is not clear, our results suggest a model in which DNA damage promotes the formation of “aberrant” forms of HR intermediates of repetitive DNA, which are recognized by QDE-3, the Neurospora homolog of the BLM/Werner helicase. Together with RPA, they recruit QDE-1 to produce aRNA and dsRNA [Lee et al. 2010]. Consistent with this model, HU treatment leads to the accumulation of rDNA-specific recombination intermediates. This model provides an explanation of why small RNAs are specifically produced at repetitive DNA loci: Only the repetitive transgenes and rDNA array provide abundant donor sequences for HR. In addition, BLM helicases have been shown to be recruited to DNA damage sites and play an important role in resolving aberrant recombination intermediates [De Muyt et al. 2012].

Repeat-associated small RNAs have been found in almost all eukaryotes. Since our report of qiRNA, DNA damage-induced small RNAs have been demonstrated in Arabidopsis, Drosophila, and mammals [Francia et al. 2012; Michalik et al. 2012; Wei et al. 2012], suggesting that DNA damage is a common trigger for small RNA production in eukaryotes. Because of the conservation of eukaryotic RNAi pathways and the fact that repetitive DNA can be a trigger for DNA damage, our results suggest that HR is also likely to be involved in small RNA production processes in other organisms.

Materials and methods

Strains and growth conditions

A wild-type strain of N. crassa (FGSCA4200) was used in this study. Mutant Neurospora strains are listed in Supplemental Table S2. The Neurospora knockout mutant strains used in this study were obtained from the Fungal Genetic Stock Center. Liquid cultures were grown in minimal medium (1× Vogel’s, 2% glucose). For liquid cultures containing QA, 10−4 M QA (pH 5.8) was added to the culture medium containing 1× Vogel’s, 0.1% glucose, and 0.17% arginine. To induce qiRNA production, histidine (1 mg/mL) or the indicated concentrations of HU were added, and cultures were collected 48 h later [Lee et al. 2009]. For race tube assays containing QA, no glucose was added to the medium.

Quelling assay, RNA Northern blot analyses, HR assay, 2D gel electrophoresis, and ChIP assay

Detailed methods for the quelling assay, RNA Northern blot analyses, HR assay, 2D gel electrophoresis, and ChIP assay are described in the Supplemental Material.

Acknowledgments

We thank Cuihong Chen and Guojun Wu for assistance in the genetic screen, and Dr. Chul-Hwan Lee for technical support. We thank Yunkun Dang, Liande Li, Qiuying Yang, Haiyan Yuan, and Annie Ye for technical assistance, and Dr. Hirokazu Inoue for providing the bar-containing plasmid for HR assay. This work was supported by grants from the National
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*Genes Dev.* 2013 27: 145-150 originally published online January 15, 2013
Access the most recent version at doi:10.1101/gad.209494.112

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